

Docket No. 44657-AAA-PCT-US/JPW/GJG/BJA

IN THE UNITED STATES PATENT AND TRADEMARK OFFICE

Applicant: Joseph R. Berger

Serial No.: 10/052,961 Group Art Unit: 1617

Filed: January 18, 2002 Examiner: S. Wang

Title : A METHOD FOR AMELIORATING MUSCLE WEAKNESS/WASTING

IN A PATIENT INFECTED WITH HUMAN IMMUNODEFICIENCY

VIRUS-TYPE 1

1185 Avenue of the Americas New York, New York 10036

Commissioner for Patents P.O. Box 1450 Alexandria, VA 22313-1450

Sir:

DECLARATION UNDER 37 C.F.R. \$1.132 OF FAITH OTTERY, M.D., PH.D., FACN

I hereby declare that:

- I am Senior Director, Medical Affairs, at Savient Pharmaceuticals, which is the owner of the above-identified application.
- 2. I am familiar with the selection and manufacture of unit dosage forms of oxandrolone. The 10mg unit dosage form of oxandrolone is approved for promoting weight gain after weight loss following chronic infection, such as HIV infection (for example, for prescribing information see Exhibit A).
- I am familiar with the specification of the above-identified application, and with each of Metcalf et al., Metabolism,

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14(1):59-66 (1965) ("Metcalf"), the description of ANAVAR® and U.S. Patent No. 5,073,380 issued to Babu et al., collectively "the cited combination of prior art."

- 4. I have read the November 15, 2007 Office Action issued in connection with the above identified application. The November 15, 2007 Office Action relies primarily on Metcalf to conclude that a 10mg unit dosage form would be obvious. I disagree with this conclusion in the November 15, 2007 Office Action.
- In my experience the nitrogen-retention ratio, as proposed 5. by Metcalf, has not been validated as a standard to be indicative of muscle mass change generally, patients specifically. Nitrogen retention as used by Metcalf is a complex interplay of a number of variables and is not necessarily indicative of muscle mass or of muscle strength. It is therefore unpredictable based on Metcalf whether the sparing" with maximum "nitrogen "optimum" for oxandrolone of 25-30mg per day would be the optimum dose for ameliorating muscle weakness or wasting in a patient such as an HIV patient.
- 6. Even if one ignores the fact that Metcalf is not directly relevant to promoting weight gain per se or weight gain specifically in HIV patients, Metcalf and the cited combination of prior art can be reasonably only interpreted to suggest a 25-30mg unit dosage form for oxandrolone.
- 7. Those in the art are aware of "pill-burden" issues as related to patient-compliance (adherence) in chronic conditions. These issues are especially important in

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patients being treated for HIV with multiple tablets. Consequently, those in the art are motivated to formulate treatments that minimize the number of tablets a patient needs to take each day. Generally, pill-burden concerns and patient-compliance issues both argue against splitting dosing into multiple tablets.

- 8. Therefore, one of skill in the art aware of the artrecognized issues of pill-burden and patient-compliance would understand Metcalf to suggest unit dosage forms of 25oxandrolone. Nothing in Metcalf. the cited combination of prior art, or the November 15, 2007 Office Action provides any rationale to split and how to split the 25-30mg optimum daily dose proposed by Metcalf for nitrogen retention. In particular, I find no reason in Metcalf and the cited combination of prior art to make a 10mg unit dose form of oxandrolone instead of a 25-30mg unit dose form or vet some other dose form.
- 9. In addition, Grunfeld et al. (1986), a copy of which is attached hereto as Exhibit B, shows that administration of a single 20mg oxandrolone tablet per day to HIV patients was statistically similar to placebo results in treating weight loss in HIV patients: "[o]nly the gain in weight at the 40mg dose of oxandrolone and the gain in BCM at the 40- and 80-mg doses of oxandrolone were greater than those in the placebo group" (see Abstract, page 304). The results in al. show that unit dosage forms Grunfeld et unpredictable effects - a 20mg unit dose of oxandrolone did not work, yet the 10mg unit dose form is approved as set forth in paragraph 2 above. A person familiar with unit dose

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formulation, therefore, could not predict from Metcalf and the cited combination of prior art which unit dosage form of oxandrolone would be effective to treat a given condition.

I hereby declare that all statements made herein of my own knowledge are true, and that all statements made on information and belief are believed to be true; and further that these statements are made with the knowledge that any such willful false statement and the like so made is punishable by fine or imprisonment, or both, under Section 1001 of Title 18 of the United States Code and that any such willful false statement may jeopardize the validity of the application or any patent issuing thereon.

1 . D Attained

Date:	5/15/08	Jaim D. Charge				v	
		Faith	D.	Ottery,	MD,	PhD	

EXHIBIT A

Applicant: Joseph R. Berger

Serial No.: 10/052,961 Filed: January 18, 2002

Exhibit A

dininistration of corticoservids, and for the relief naintain normal weight, to offset the protein of the bone pain frequently accompanying catabolism associated with prolonged ostoporosis (Scr DOSAGE AND ADMINISTRATION).

SAVIENT PHARMACEUTICALS, INC.

Daandrin oral table to contain 2.5 mg or 10 mg

Oxandrokane is 17\$-hydraxy-17a-methyl-2-

of the anabolic steroid exandralone. DESCRIPTION

oxa-50-androstan-3-one with the following

structural formula:

Oxandria@(oxandrolone tabletx, USP)Ciii

Act of 1990 and has been assigned to Schedule III substance under the Anabolic Steroids Control DRUG ABUSE AND DEPENDENCE Oxandrolone is classified as a controlled

CONTRAINDICATIONS

Carcinoma of the breast in females with Known or suspected carcinoma of the steroids may stimulate osteolytic bone sypercalcemia (androgenic anabolic prostate or the male breast. ر.

nactive ingredients include cornstanch, factore,

magnesium stearate, and hydroxypropyl

methylcellulose.

has been shown to cause embryotoxicity, masculinization of the fetus. Oxandrin nasculinization of female animal Pregnancy, because of possible ctotoxicity, infertility, and resorption). -i

testosterone. Certain chaical effects and adverse

Anabolic steroids are synthetic derivatives of

CLINICAL PHARMACOLOGY

reactions demonstrate the androgense properties of this class of drugs. Complete dissociation of herefore similar to those of male sex hormones

anabolic and androgenic effects has not been achieved. The actions of anabolic steroids are

Nephrosis, the nephrotic phase of he human dose.

offspring when given in doses 9 times

nephrits. Hypercalcenia.

disturbances of growth and sexual development if

with the possibility of causing serious

given to young children. Anabolic steroids suppress the gonadotropic functions of the vitalizary and may exert a direct effect upon the

indrogens, endogenous testosterone release is

During exogenous administration of anabolic

inhibited through inhibition of pituliary lucinizing hormone (LH). At large doses, spermanogenesis may be suppressed through feedback inhibition of pituliary follicle.

SOMETIMES PRESENT WITH MINIMAL PELIOSIS HEPATIS, A CONDITION IN SPLENIC TISSUE IS REPLACED WITH REPORTED IN PATIENTS RECEIVING ASSOCIATED WITH LIVER FAILURE. THEY ARE OFTEN NOT RECOGNIZED WITHDRAWAL OF DRUG USUALLY ANDROGENIC ANABOLIC STEROID BLOOD-FILLED CYSTS, HAS BEEN HEPATIC DYSFUNCTION, BUT AT OTHER TIMES THEY HAVE BEEN UNTIL LIFE-THREATENING LIVER FAILURE OR INTRA-ABDOMINAL WHICH LIVER AND SOMETIMES INERAPY. THESE CYSTS ARE HEMORRHAGE DEVELOPS.

Anabolic storoids have been reported to increase

timulating hormone (FSH).

ow-density lipoproteins and decrease high-

lensity lipoproteins. These levels nevert to

tormel on discontinuation of treatment.

n a single dose pharmacokinetic study of climination half-life was 13.3 bours. In a

Oxandrin is elderly subjects, the mean

RESULTS IN COMPLETE DISAPPEARANCE OF LESIONS.

younger volunteers, the mean climination half-life found for time to peak, peak plasma concentration

setween younger and elderly voluntoers were or AUC after a single dose of Oxandrin. The

was 10.4 hours. No significant differences

previous single dose pharmacokinetic study in

TUMOR, HOWEVER, HEPATIC TUMORS ANDROGEN-DEPENDENT, BUT FATAL MALIGNANT TUMORS HAVE BEEN REPORTED. WITHDRAWAL OF DRUG CESSATION OF PROGRESSION OF THE **OFTEN RESULTS IN REGRESSION OR** ASSOCIATED WITH ANDROGENS OR SILENT UNTIL LIFE-THREATENING ANABOLIC STEROIDS ARE MUCH REPORTED. MOST OFTEN THESE MORE VASCULAR THAN OTHER HEPATIC TUMORS AND MAY BE LIVER CELL TUMORS ARE ALSO TUMORS ARE BENIGN AND

conclusion between plasma level and therapeutic

:feet has not been defined.

ANDROGENS OR ANABOLIC STEROIDS. BE VERY MARKED AND COULD HAVE ATHEROSCLEROSIS AND CORONARY LIPOPROTEINS. THE CHANGES MAY A SERIOUS IMPACT ON THE RISK OF ATHEROSCLEROSIS ARE SEEN IN LIPOPROTEINS AND SOMETIMES DECREASED HIGH-DENSITY INCREASED LOW-DENSITY WITH INCREASED RISK OF PATIENTS TREATED WITH THESE CHANGES INCLUDE ARTERY DISEASE.

with joundice appears or if liver function tests Cholestatic hepatitis and jaundice may occur become abnormal, oxandrolone should be discontinued and the etiology should be relatively low dose. If cholestatic hepatitis with 17-alpha-alkylated androgens at a letermined. Drug-induced jaundice is reversible when the medication is

patients with breast cancer, anabolic steroid therapy may cause hypercalcemia by stimulating osteolysis. Oxandrolone therapy hould be discontinued if hypercalcenia

sepair disease. Concomiant administration sations with pre-existing cardiac, renal, or of adrenal contical steroid or ACTH may Edema with or without congestive bear failure may be a serious complication in acrease the odenta.

height. The younger the child, the greater the risk of compromising final mature height. n children, androgen thorapy may accelerate adverse effect results in compromised adult monitored by assessing bone age of the left wrist and hand every 6 months (See bone metaration without producing compensatory gain in linear growth. This The effect on bone maturation should be PRECAUTIONS: Laboratory Tests).

unabolic steroids may be at an increased risk for the development of prostatic hyportrophy Ceristric patients treated with androgenie and prostatic carcinome. ANABOLIC STEROIDS HAVE NOT BEEN SHOWN TO ENHANCE ATHLETIC

AND ADMINISTRATION.

(PT). When exandreione is prescribed to patients being treated with warfarin, dones warfaria may result in anexpectedly large increases in the INR or prothrombin time Concurrent desing of exandrolene with PRECAUTIONS

level and chminish the risk of potentially serious bleeding (See PRECAUTIONS: Drug Lateractions).

sene, choromogaly). Discuntinuation of drug virilization (deepening of the voice, hirsuism, virilism is necessary to prevent irreversible virilization. Some virilizing changes in sevented by concomitant use of estroyens vocaca are irreversible even after prompt Women should be observed for signs of acrapy at the time of evidence of maid Menserual irregularities may also occur. iscontinuance of therapy and are not

loating factors II, V, VII, and X, and an increase tasbolic steroids may cause suppression of a prothrombis time.

The physician should instruct patients to report The physician should instruct patients to report my of the following side effects of androgens: Makes. Too frequent or persistent erections of he penis, appearance or aggravation of aene. til pariemer. Nausca, vomiting, changes in innediately any use of warfarin and any emales: Houseness, w.mc, changes in nenstrust periods, or more facial hair. kis color, or ankle swelling. nformation for patients:

imiter between the two age groups although the Oxandrin, at daily doses of 5 mg bid, and 10 mg everage duration of treatment from 68.5 days to ge. No significant differences in efficacy were descend between the 5 mg bid and 10 mg bid inderlying medical conditions. The maximum evented as increased half-life when compared cessitivity to drug-induced fluid retention and avolving a total of 339 paintats with different idealy patients (2 65 years of age) received timilar in those 2 65 and those < 65 years of centiivity to fluid resention and increases in furstion of incannent was 4 months with the bally doses. The adverse event profiles wert iderly, particularly is women, had a greater M.7 days across the studios. A total of 172 Duandria treatment. Mean weight gain was harmacokinetic study in elderly volunteers recommended in the elderly (see DOSAGE ransaminase elevations, a lower dose is PHARMACOLOGY) Based on greater o younger volunteers. (see CLINICAL aid, was evaluated in four clinical trads separic transminance. A single dose

should have frequent desermination of unite and Women with discerningted breast carcinoma serum calcium levels during the course of Laboratory Tests:

scrapy. (See WARNINGS).

THAT ARE KNOWN TO BE ASSOCIATED

of warfaria may need to be decreased

INTRA-ABDOMINAL HEMORRHAGE

rauma, and in some patients who without definite

Oxandrin is indicated as adjunctive thempy to promote weight gain after weight loss following extensive surgery, chronic infections, or severe

INDICATIONS AND USAGE

overuse of the reparatoxicity assectated with the use of 17-slpha-allylated androgens, liver function tests should be obtained periodically.

bone age should be made during measurest of children to determine the rate of bone maturation Periodic (every 6 months) x-ray examinations of and the effects of anthrogen therapy on the epiphyseal centers.

Androgenic unabolic steroids have been reported cardiovascular disease of who are at risk for cardiovascular disease. Serum determination of decrease high-density inapproteins. Therefore, castion is required when administering these lipid levels should be performed periodically to increase low-density importoients and agends to patients with a history of and therapy adjusted accordingly.

Hemoglobin and hematocrit should be checked periodically for polycythenia in patients who are receiving high doses of anabolic steroids.

Drug interactions

oral anticoagulants. Dosage of the anticoagulant may have to be decreased in order to maintain Anabulic steroids may increase sensilivity to monitaring, especially when anabolic steroids desired producembin time. Patients receiving oral anticoagulant therapy require close are started or stopped.

from 6, 13 mg/day to 1, 13 mg/day (approximately 10, 1855; reaction of warfarin does, was necessary to manioning a unger INR of 1.5. When connectedore therapy is indinated in a patient afterly precising transmit with warfarin, the INR or proformable time (PT) thould be invasioned closesty and the done of warfarin balf-life from 26 to 48 hours and AUC tingival bleeding (1/15) were also observed. A from 4.55 to 12.08 ng*hr/nM.; similar increases in N-warfarin half-life and AUC were also furthermore, in patients receiving both drugs, telected. Microscopic hemannia (9/15) and 5.5-fold decrease in the mean warfaria dose ovandrolone, given as 5 or 10 mg bid in 15 warferin adjusted as necessary until a stable healthy subjects concurrently treated with marfaria, resulted in a mean increase in Scareful monitoring of the INR or PT, and Werfarin: A muhidose study of larges INR or P.T has been achieved.

In patients with edense, concomitant administration with adrenal cortical steroids or ACTH may increase the oderna. Adrenal steroids or ACTH:

thyrozine-binding globulin, resulting in dereuted total farmus levels and increased test of the markets and increased treats uptake of 1 and T. Free laying burnoas berels remain unchanged. In addition, a decrease in PBI and radioactive todine uptate. Anabalic steroids may decrease levels of Srug/Laboratory test interactions: may occur.

Sarcinegenesis, mutagenesis, impairment of ertility

unimals for carcinogenic or mutagenic effects. In 2-year chronic oral ras studies, a dose-related organ weights (lestes, prostate, seminal vesicles, Oxendrolope has not been tested in laboratory overies, uterus, adrenals, and pituitary) were reduction of spermatogenesis and decreased POWE

derna data:

WARNINGS). Withdrawal of the drugs did not liver cell tumors have been reported in patients receiving long-term therapy with sackrogenic cad to regression of the tumors in all cases. mabolic servids in high doses (See

trabolic steroids may be at an increased risk for he development of prostatic hypertrophy and Deniatric patients treated with androgenic Profitic carcinoma.

Category X (See CONTRAINDICATIONS). repassor: Teratogenic effects-Pregnancy

Fluid and electrolytes: Edema, retention of

terum electrolytes (sodium chloride,

potential of scrious adverse reactions in nursing afants from oxandrolone, a decision should be t is not known whether anabolic steruids are made whether to discontinue nursing or to discontinue the drug, taking into account the acresed in human milk. Because of the reportance of the drug to the mother. iursing mothers:

solcrance (See PRECAUTIONS: Laboratory

increased creatinine excretion,

norcased serum levels of creatinine

Metabolic/Endocrine: Decreased glucose

sotssium, phosphate, calcium).

he fetus. Inhibition of gonadutupin scretton

busphokinase (CPK). Masculinization of

herefore, thorapy abould be monitored by x-ray studies at 6-month intervals in order to avoid the risk of compromising adult height. Androgenic naturation more rapidly than linear growth in unabolic agents may accolurate epiphysen! children and the effect may continue for 6 mouths after the drug has been stopped.

or COPD exacerbation and fluid netention. ADVERSE REACTIONS

noophams and policius hepatis with long-term thorapy (See WARNINGS). Reversible changes in liver function tests also occur including increased beconsult ophatises (BSP) retriction, changes in alla-line phosphatase and uninousnsferase (AST, SGOT) and alsaine The following adverse reactions have been Hepatic: Cholestalic jaundice with, rarely, repatic necruais and death. Hepatocellular associated with use of anabolic steroids: nortaxes in senum bilirubin, aspartate uninotransferase (ALT, SGPT) " males.

100 (NDC 54396-111-11) Pastpubertal: Inhibition of testicular function. testicular amophy and oligospermia, impotence, Cliand margement, mensurusi irregularities. chronic priapism, epididymitis, and bladder CMS: Habituation, excitation, insormia, depression, and changes in libido. Heranologic: Bleeding in patients on

Tribulary In females R, only

Savient Phermacrusicals, Inc. by:

Luryna: Despening of the voice in females. Hair: Hirmism and male partern baldness

concomitant aral anticoagulant therapy.

Breast, Gynecomagia,

NSM Pharmaccuticals, Inc. steamile, NC 27834

few York, NY 10017

Address medical inquires to: Savient Pharmacenéesis, Inc. East Brunswick, NJ 08816 166-692-6374 One Towar Center ourteenth Floor

\$2005, Saviest Phermaceuticals, Inc. Printed in USA

(oxandrologe tablets, USP)CIII OXANDRING

SAVIENT

The oral LD to of oxandrolone in mice and dogs

intidote is known, but gastric lavage may be

DOSAGE AND ADMINISTRATION

Thempy with anabolic steroids is

djunctive to and not a replacement for

is greater than 5,000 mg/kg. No specific

No symptoms or signs associated with overdosage have been reported. It is possible

OVERDOSAGE

that sodium and water retention may occur.

conventional therapy. The duration of therapy with Oxandrol (oxandrolene) who are aware of the effects on bone maturation See WARNINGS).

Oral hypoglycemic agents: Oxunhylone may inhibit the metabolism of oral

cautiously in children and only by specialists

mabolic steroid therapy should be used very

are recommended when the exandrolone dose is

changed or discontinued. Patients should be

closely axonitored for signs and symptoms of

occult bleeding.

adjustment of the warfarin dosage if indicated

stanchadilators should be monitored closely Patients with moderate to severe COPD or COPD patients who are unresponsive to

patient and the possible appearance of alverse reactions. Therapy should be

will depend on the response of the

fedula: The response of impividuals to anabolis

servids varies. The daily adult desage is 2.5 mg This may be repeated intermittently as indicated Children: For children the total daily datage of issued response may be achieved with as link is 2.5 mg or as much as 20 mg daily. A course Oxandrin is 50.1 mg per kilogenen body weight of therapy of 2 to 4 works is usually adequate. or 50.045 mg per pound of budy weight. This to 20 mg given in 2 to 4 divided dozen. The nay be repeated intermittently as indicated.

Geriatric Use: Recognicaded dose for geriatric patients is 5 mg bid.

Prepuberul: Phaltic enlargement and increased

requency or persistence of erections.

HOW SUPPLIED

ide of the scoreline on the other side; bonles of scored with BTG on one side and "11" on each wher side; bottles of 60 (NDC 54396-110-60). white, with BTO on one side and "10" on the Oxandria 2.5 mg tablets are oval, white, and Oxandrin 10 mg tablets are capsule shaped.

Issued: Lanuary, 2006

Manufactured for

Meletal: Promature closure of epiphyses

in children (See PRECAUTIONS:

"calatric ace).

Skir: Acne (especially in females and

in females.



Applicant: Joseph R. Berger

Serial No.: 10/052,961 Filed: January 18, 2002

Exhibit B

CLINICAL SCIENCE

Oxandrolone in the Treatment of HIV-Associated Weight Loss in Men

A Randomized, Double-Blind, Placebo-Controlled Study

Carl Grunfeld, MD. PhD,* Donald P. Kotler, MD,† Adrian Dobs, MD,‡ Marshall Glesby, MD,§ and Shalender Bhasin, MD, // for the Oxandrolone Study Group

Objective: To evaluate the efficacy and safety of oxandrolone in promoting body weight and body cell mass (BCM) gain in HIVassociated weight loss.

Methods: Randomized, double-blind, placebo-controlled trial. Two hundred sixty-two HIV-infected men with documented 10% to 20% weight loss or body mass index ≤20 kg/m2 were randomized to placebo or to 20, 40, or 80 mg of oxandrolone daily. After 12 weeks, subjects were allowed to receive open-label oxandrolone at a dose of 20 mg for another 12 weeks.

Results: Body weight increased in all groups, including the group receiving placebo, during the double-blind phase (1.1 ± 2.7, 1.8 ± 3.9, 2.8 ± 3.3, and 2.3 ± 2.9 kg in placebo and 20-, 40-, and 80-mg oxandrolone groups, respectively; all P < 0.014 vs. baseline). BCM increased from baseline in all groups (0.45 ± 1.7, 0.91 ± 2.2, 1.5 ± 2.5, and 1.8 ± 1.8 kg in placebo and 20-, 40-, and 80-mg oxandrolone groups, respectively). At 12 weeks, only the gain in weight at the 40-mg dose of oxandrolone and the gain in BCM at the 40- and 80-mg doses of exandrolone were greater than those in the placebo group, however. Oxandrolone treatment was associated with significant suppression of sex hormone-binding globulin, luteinizing hormone, follicle-stimulating hormone, and total and free testosterone levels. Treatment was generally well tolerated but accompanied by significant increases in transaminases and lowdensity lipoprotein as well as decreases in high-density lipoprotein. Cenclusion: Oxandrolone administration is effective in promoting dose-dependent gains in body weight and BCM in HIV-infected men with weight loss.

Key Words: wasting syndrome, cachexia, anabolic therapy, body composition, anabolic steroid, lean body mass, fat toxicity, liver function, lipoproteins, atherosclerosis

(J Acquir Immune Defic Syndr 2006;41:304-314)

Ithough the prevalence of weight loss in HIV-infected Annual though the prevalence of worght spread use of antiretroviral drug therapy, weight loss continues to be a significant problem, affecting 31% of patients during the course of their illness. 1-4 In Africa and Asia, where most HIV-infected patients reside, weight loss is a major presenting feature of AIDS.5 Weight loss and, in particular, loss of body cell mass (BCM) are independent risk factors for death in patients with HIV infection, even when the CD4 cell count and history of complications are taken into account.⁵⁻¹¹ Furthermore, loss of weight, lean body mass, and BCM are accompanied by decreased function, worsening quality of life (QOL), and increasing hospitalization rates. 12-13

Early studies suggested that BCM was preferentially lost and fat spared in men with HIV-associated wasting. 16 Whereas some subsequent studies had similar findings, other studies in men and women found more significant loss of fat. 18-21 An explanation for these discrepancies is that subjects who had a low percentage of fat when first studied lost predominantly lean body mass, whereas those who started with higher percentage fat lost predominantly fat.²⁰

Nutritional therapy and appetite stimulants can promote weight gain in patients with HIV-associated wasting, 22-25 The predominant gain is in body fat, however. Although increased energy stores may reduce loss of BCM in future episodes of weight loss, 18 body fat stores do not correlate with survival. 6.8.9 In contrast, anabolic therapy with growth hormone (rhGH) has the potential of inducing gain of lean body mass; however, rhGH therapy also induces loss of fat reserves.²⁶⁻²⁶

Testosterone supplementation increases fat-free mass and muscle strength in HIV-infected men with mild to moderate weight loss. 30-37 Androgenic steroids promote a positive nitrogen balance and weight gain (or amelioration of weight loss) in other catabolic illnesses, including acute alcoholic hepatitis, cancer, end-stage renal disease, and burns. 18-51 In studies of small numbers of patients with HIV-associated wasting, orally administered androgens, such as oxandrolone and oxymetholone, and the parenterally administered androgen nandrolone decanoate have induced significant weight gain. 43-51 Given the potential advantage of an orally administered anabolic therapy, such as oxandrolone, we undertook a double-blind, placebo-controlled, randomized trial of graded doses of oxandrolone in HIV-infected subjects with weight loss, testing its effects on weight gain, body composition, total work capacity, health-related QOL, and safety.

Received for publication August 3, 2005; accepted November 10, 2005.

Received for publication rugust 5, 2007, excepts a vice from the *University of California, San Francisco, and the Department of Veterans Affairs Medical Center, San Francisco, CA; 15t. Lukes-Roosevelt Medical Center, Columbia University School of Medicine, New York, NY: \$Johns Hopkins School of Medicine, Baltimore, MD; \$Community Research Initiative on AIDS, New York, NY and ||Charles Drew University of Medicine and Science, Los Angeles, CA.
Grant support provided by Biotechnology General (now Savient Pharmaceu-

Reprints: Carl Grunfeld, Metabolism section (111F), Department of Veteran Affairs Medical Center, 4150 Clement Street, San Francisco, CA 94121 (e-mail: grunfid@itsa.ucsf.edu).

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METHODS

Signed informed consent was obtained from each patient before entry under protocols approved by the institutional review board at each participating center. This was a randomized, placebo-controlled, parallel-group, double-blind, multisite clinical trial conducted at 25 sites between September 25, 1996 and July 20, 1998.

Participants

Eligible subjects were HIV-infected men ≥18 years of age who had 10% to 20% unintentional weight loss from premorbid weight documented in medical records or a body mass index (BMI) ≤20 kg/m², a Karnoſsky Performance Scale score >60%, a liſe expectancy of >6 months, and the ability to consume a normal well-balanced diet at entry as assessed by a dietitian. Therapy with antiretroviral medication was not required; however, subjects on antiretroviral therapy had to be on a stable regimen for more than 6 weeks at the time of entry.

Exclusion criteria included any opportunistic infection within 60 days of enrollment; loss of >5% body weight in the previous 30 days; chronic fever >101°F with a frequency ≥3 days per week for at least 2 weeks in the previous 30 days; aspartate aminotransferase (AST), alanine aminotransferase (ALT), or alkaline phosphatase levels greater than 5 times the upper limit of normal and/or bilirubin level ≥2.0 mg/dL within 2 weeks; serum creatinine >2.0 mg/dL; known impaired digestive or absorptive function; chronic uncontrolled diarrhes (>3 liquid stools per day at least 4 days per week for >2 weeks); current treatment with anticoagulants or oral hypoglycemic agents; or treatment with appetite stimulants, weight-promoting agents, anabolic steroids, or testosterone in the previous 4 weeks. This dose-ranging study did not include women, because it was not known whether doses of this magnitude would cause significant virilization in women.

Treatment Assignment and Randomization

Subjects were assigned in concealed randomization (1:1:1:1) balanced at each center to placebo (4 tablets) or to 20 mg/d of oxandrolone (1 20-mg tablet of oxandrolone and 3 placebos), 40 mg/d (2 20-mg tablets of oxandrolone and 2 placebos), or 80 mg/d of oxandrolone (4 20-mg oxandrolone tablets) provided by Savient Pharmaceuticals (East Brunswick, NJ [formerly Bio-Technology General, Corporation]). Investigators and patients were blinded to treatment assignment during the initial 12 weeks. After 12 weeks, all subjects who wished to continue were placed on 20 mg of oxandrolone in an open-label continuation.

Subject Accountability

Two hundred sixty-two patients were randomized and included in the intent-to-treat analysis (placebo [n=65], 20 mg of oxandrolone [n=64], 40 mg of oxandrolone [n=65], and 80 mg of oxandrolone [n=68]). Of these, 195 subjects completed the double-blind phase and 193 completed the open-label phase. Of the 67 subjects who discontinued treatment during the double-blind phase, 12 were in the placebo group, 18 were in the 20-mg oxandrolone group,

18 were in the 40-mg oxandrolone group, and 19 were in the 80-mg oxandrolone group. Reasons for discontinuations included adverse experience, ²⁰ death, ⁶ intercurrent medical problem or disease-related complication, ² subject relocation or voluntary patient withdrawal, ²¹ and noncompliance. ¹⁸

Assessments

Measurements were made at baseline and at 2, 4, 8, and 12 weeks in the double-blind placebo phase and at weeks 14, 18, and 24 in the open-label study. The primary outcome was change in body weight measured at each time point under standardized conditions (at the same time, preferably in the morning, wearing only underwear and socks) on a single balance-type scale that had been recently calibrated by a state agent or third-party source. Other outcomes included measurement of fat and BCM by bioelectrical impedance analysis (BIA) (RJL Systems). Because changes in hydration status and technical aspects of performance affect BIA, data were excluded if the change in BCM was >2.5 times the change in weight or the change in BCM was >7.5 kg in a subject; 18 subjects were thus excluded from the body composition analysis because of quality control problems with BIA measurements (6 from placebo group, 3 from 20-mg oxandrolone group, 6 from 40-mg oxandrolone group, and 3 from 80-mg exandrolone group). Health-related QOL was measured by the Medical Outcomes Study (MOS) HIV health survey.⁵² Treadmill tests were performed at centers with treadmill capability on day I and at weeks 4 and 12. Changes in physical capacity were assessed by changes in total workload from the treadmill tests. Nineteen percent of subjects had treadmill tests performed at week 12. Total workload is defined as $\Sigma[\text{speed (m/min)}]$ [% grade/100] [time in minutes on treadmill].

Safety assessments, including HIV RNA levels by reverse transcriptase polymerase chain reaction (RT-PCR), CD4 T-lymphocyte counts, complete blood cell counts, and blood chemistry, were measured at Covance Laboratories.

Serum luteinizing hormone (LH), follicle-stimulating hormone (FSH), and sex hormone-binding globulin (SHBG) as well as testosterone levels were measured by 2-site-directed immunofluorometric assays (Delfia-Wallac, Gaithersburg, MD), with sensitivities of 0.05 U/L, 0.15 U/L, and 6.25 nmol/L, respectively, as described previously. ^{53,54} The intra- and interassay coefficients of variation were 10.7% and 13.0% for LH, 3.2% and 11.3% for FSH, and 10.0% and 10.2% for SHBG, respectively. The cross-reactivity of free a-subunit and other pituitary hormones in the LH and FSH assays was <1%.

Serum total testosterone levels were measured using a radioimmunoassay (RIA) with an iodinated testosterone tracer^{4.35} that has been validated against liquid chromatography-mass spectrometry tandem mass spectrometry. This assay has a sensitivity of 0.44 ng/dL and intra- and interassay coefficients of variation of 8.2% and 13.2%, respectively. Free testosterone levels were measured by a sensitive equilibrium dialysis method.^{54.35} optimized to measure low concentrations with accuracy. Two hundred microliters of serum in the inner compartment was dialyzed against 2.4 mL of dialysis buffer that approximates the composition of a

protein-free ultrafiltrate of human scrum. Dialysis was performed overnight for 16 hours at 37°C. Testosterone concentration in the dialysate was measured by RIA using 1231-labeled testosterone. The sensitivity of the free testosterone assay is 0.6 pg/mL (2.0 pmol/L), with intra- and interassay coefficients of variation of 4.2% and 12.3%, respectively.

Total and free testosterone concentrations were not consistently changed during oxandrolone treatment despite suppression of LH concentrations, suggesting that oxandrolone or one of its metabolites might have cross-reacted in the testosterone assays. Therefore, we established a chromatographic system to separate testosterone from oxandrolone before RIA. Serum samples were extracted using ethyl acetate and hexane (3:2 vol/vol) and subjected to chromatography on celite columns equilibrated in isooctane. Lipemic samples were clarified by centrifugation before extraction. Testosterone was cluted by washing columns with 10% ethyl acetate in isooctane. In preliminary experiments, we demonstrated that >90% of ¹⁴C-testosterone eluted with 10% isooctane, whereas >90% of 3H-oxandrolone eluted with ≥15% iso-octane. Less than 5% of 14C-oxandrolone cluted with 10% isooctane; conversely, less than 5% of testosterone eluted at isooctane concentrations ≥15%. Eluates were dried under nitrogen and taken up in assay buffer. Recovery of known amounts of testosterone added to charcoal-stripped serum samples during extraction and celite chromatography was consistently better than 80%. Therefore, values were not corrected for losses during chromatography.

At each visit, intercurrent illnesses, symptoms, and additional medicines were recorded. Compliance was assessed by pill count.

Statistics

Results are presented as mean ± standard deviation (SD). Primary efficacy end points were changes in body weight and body composition from baseline. Based on preliminary data, the study was designed to detect a 2.0-kg (SD = 3.5 kg) increase in oxandrolone-treated patients compared with patients treated with placebo with a power of 80%, under the presumption that those on placebo would, on average, lose weight during the course of the study. The

study was not powered to detect significant changes in secondary end points, such as quality of life. Analysis of variance (ANOVA) and the Dunnett t test were used to analyze primary and secondary efficacy parameters. To control the overall type I error rate of 0.05 for the multiple comparisons. Bonferroni inequality was used; treatment differences were considered significant if the significance level for that comparison was <0.017 instead of 0.05. Withintreatment changes from baseline were tested using a 1-way t test. The number of patients with adverse events and discontinuations was compared using the Fisher exact test. The prevalence of World Health Organization (WHO) grade III and IV toxicities was compared with placebo using the x2 test, with differences across dosages analyzed by the Cochran-Armitage trend test. Demographic and disease history variables at baseline were compared between treatment groups using ANOVA. The effect of race was tested using the x2 test.

RESULTS

Subject Characteristics

Baseline characteristics of the subjects were not significantly different among treatment groups (Table 1). Seventy percent of participants were white, 17% were African American, 11% were Hispanic, and 2% were other. Weight loss before entry averaged 16.4% ± 8.0% from baseline.

Body Weight and Composition

In subjects who were evaluated at baseline and received drug, weight increased progressively in all groups, including the placebo group, during the study (Fig. 1A). A significant increase occurred as early as 2 weeks after baseline for each group, including the placebo group. On an intent-to-treat basis at 12 weeks or at last measurement during the double-blind placebo phase, for the 258 subjects with baseline weight (Table 2), there was a gain of 1.1 \pm 2.7 kg on placebo, 1.8 \pm 3.9 kg on 20 mg of oxandrolone, 2.8 \pm 3.3 kg on 40 mg of oxandrolone, and 2.3 \pm 2.9 kg on 80 mg of oxandrolone (all P < 0.014 vs. baseline). Weight gain at 2, 4, 8, and 12 weeks on the 40-mg dose of oxandrolone was statistically different from weight gain on placebo (P = 0.0040 vs. placebo at

TARLE 1 Baseline Characteristics of the Participants (N = 262)

	Placebe (a = 65)	Oxendrolone				
		20 mg (n = 64)	40 mg (n = 65)	88 mg (n = 68		
Age (y)	41.7 ± 8.4	41.1 ± 9.0	40.1 ± 7.5	39.5 ± 7.5		
Height (in)	69.8 ± 3.0	69.8 ± 3.2	69.0 ± 3.0	70.0 ± 2.9		
Weight (kg)	66.6 ± 9.9	65.9 ± 9.6	65.0 ± 11.1	65.4 ± 8.7		
BMI (kg/m²)	20.7 ± 2.7	21.0 ± 2.7	21.1 ± 3.2	20.6 ± 2.4		
Weight loss* (% from baseline)	17.7 ± 6.7	15.8 ± 6.1	15.0 ± 7.2	16.9 ± 11.1		
CD4* lymphocytes × 10 ^h /L	225 ± 188	226 ± 223	261 ± 211	252 ± 191		
IIIV PCR (log/ml.)	5.31 ± 5.76	5.19 ± 5.58	5.19 ± 5.58	5.09 ± 5.59		

^{&#}x27;n = 59 for placebo, n = 62 for 20 mg of exandrolone, n = 60 for 40 mg of exandrolone, and n = 60 for 80 mg of exandrolone; some subjects met entry criteria based on having a

BMI ≤ 20 and did not have weight loss recorded.

Values are mean ± SD. There were no significant differences between the groups.

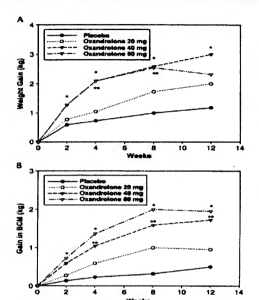


FIGURE 1. A, Average body weight gain during the 12-week treatment period. Mean change in weight from baseline at each time point is shown. Data are mean ± 5D. *P (40 mg of oxandrolone vs. placebo) < 0.017, **P (80 mg of oxandrolone vs. placebo) < 0.017, B, Average change in BCM during the 12-week treatment period. BCM was measured by BIA. Data are mean ± 5D. *P (40 mg of oxandrolone vs. placebo) < 0.017, **P (80 mg of oxandrolone vs. placebo) < 0.017.

12 weeks). The difference in weight gain between the 80-mg oxandrolone group and the placebo group was significant at 4 and 8 weeks but not at 2 or 12 weeks (P = 0.045 at 12 weeks, which did not meet the multiple comparisons criterion). There was no significant effect of performance site.

Body composition was measured by using BIA. Thirty patients at 4 centers underwent dual energy x-ray absorptiometry (DEXA) measurements to validate body composition measurements by BIA. The correlation between the measurements of fat-free mass by the 2 methods was 0.937 (P < 001).

BCM increased progressively and significantly in all groups (see Fig. 1B). At 12 weeks or last visit on an intent-to-treat basis, the increase in BCM was 0.45 ± 1.7 kg on placebo, 0.91 ± 2.2 kg on 20 mg of oxandrolone, 1.5 ± 2.5 kg on 40 mg of oxandrolone, and 1.8 ± 1.8 kg on 80 mg of oxandrolone (see Table 2). The increase in BCM on the

TABLE 2. Change in Body Weight and Composition at Week 12 (Intent-to-treat)

		Oxandrolone			
	Placebo	20 mg	40 mg	80 mg	
Weight (kg)	1.1 ± 2.7	1.8 ± 3.9	2.8 ± 3.3*	2.3 ± 2.9	
n	64	63	64	67	
BCM (kg)	0.45 ± 1.7	0.91 ± 2.2	1.5 ± 2.5†	1.8 ± 1.82	
n	62	61	59	64	
Intracellular water (L)	0.4 ± 1.6	0.8 ± 2.0	1.4 ± 2.3†	1.7 ± 1.62	
n	62	61	59	64	
Extracellular water (L)	0.3 ± 1.5	0.4 ± 2.9	0.2 ± 1.3	-0.2 ± 1.5	
n	62	61	59	64	
Body fat (kg)	0.3 ± 1.6	0.4 ± 2.2	1.0 ± 2.4%	0.6 ± 1.8	
n	62	61	59	64	

^{*}P = 0.004 vs. placebo

40-mg and 80-mg doses at 12 weeks was significantly greater than that on placebo (P=0.0049 and P=0.0002, respectively). Similar results were obtained when intracellular water was analyzed by BIA (see Table 2). In contrast, there were no significant changes in extracellular water in any group (see Table 2). There was also a trend to gain body fat on the 40-mg dose, but this did not reach statistical significance using the multiple comparisons criteria.

The entry criteria included 10% to 20% of unintentional loss of weight or a BMI ≤20 kg/m². These criteria allowed patients who were over their ideal body weight or even obese at baseline to enter the study if they had lost 10% to 20% of their body weight. Five subjects were obese (>120% ideal body weight), with the highest weight at entry being 107 kg. Twenty-four percent of the patients had a BMI >22.5 kg/m². Therefore, we performed post hoc analysis evaluating changes in body weight and composition in subjects with a BMI ≤22.5 kg/m² on an intent-to-treat basis at 12 weeks or last measurement. Their weight increase over baseline at 12 weeks was 0.8 ± 2.7 kg on placebo, 2.7 ± 4.0 kg on 20 mg of oxandrolone, 2.9 ± 2.6 kg on 40 mg of oxandrolone, and 2.5 ± 2.8 kg on 80 mg of oxandrolone (all significantly increased over baseline). Compared with placebo, subjects receiving 20 mg, 40 mg, or 80 mg of oxandrolone had significantly higher weights at week 12 (P = 0.0026, P = 0.0005, and P = 0.00050.0041, respectively). Similar changes were found for BCM, where the increases at 12 weeks over baseline were 0.2 ± 1.5 kg in the placebo group, 1.1 ± 2.1 in the 20-mg oxandrolone group; 1.8 ± 1.5 kg in the 40-mg oxandrolone group, and 2.0 ± 1.7 kg in the 80-mg oxandrolone group. Compared with placebo, subjects receiving 20, 40, or 80 mg of oxandrolone had a significantly higher BCM at week 12 (P = 0.0122, P < 0.0001, and P < 0.0001, respectively).

Functional Outcomes

No significant differences were seen in MOS HIV health surveys for any treatment group. There was no significant change from baseline in total work output in the

tP = 0.0049 vs. placebo tP = 0.0002 vs. placebo

IP = 0.0450 vs. placebo.

subset of subjects who underwent treadmill testing in any treatment group. There was no correlation between change in weight and QOL score or total work output.

Safety

Neither HIV RNA by RT-PCR nor CD4 * lymphocyte count was significantly affected by oxandrolone (Table 3). There were no significant changes in hemoglobin and white blood cell counts. However, there was a dose-dependent increase in platelet count (P < 0.017 for all doses of oxandrolone vs. placebo). There were small but significant increases in levels of creatinine and creatine kinase but not in blood urea nitrogen (BUN) in the oxandrolone groups compared with the placebo group.

Serum albumin, total protein, bilirubin, alkaline phosphatase, lactate dehydrogenase (LDH), and gamma-glutamyl transferase (GGT) levels were not significantly changed (see Table 3). However, there were dose-dependent increases in AST and ALT appearing by the first 4 to 8 weeks of therapy. The increase in AST was significant at the 80-mg dose compared with baseline, whereas the increase in ALT was significant.

cant at the 40-mg and 80-mg doses. Furthermore, there was a dose-related increase in the incidence of WHO grade III and IV liver toxicity for ALT and AST with increasing dose of oxandrolone (Table 4). For AST, WHO grade III and IV toxicity occurred in 2 of 61 subjects on placebo, 2 of 60 on 20 mg of oxandrolone, 6 of 61 on 40 mg of oxandrolone, and 9 of 61 on 30 mg of oxandrolone. For ALT, WHO grade III and IV toxicity occurred in 1 of 61 subjects on placebo, 3 of 60 on 20 mg of oxandrolone, 7 of 61 on 40 mg of oxandrolone, and 9 of 61 on 80 mg of oxandrolone (for trend, P = 0.0047). Three subjects receiving the 40-mg dose and 4 subjects receiving the 80-mg dose were discontinued from the drug because of laboratory abnormalities.

Glucose, triglyceride, and total cholesterol levels in patients receiving oxandrolone were not significantly different from those receiving placebo (see Table 3). There was a significant decrease in uric acid and plasma high-density lipoprotein (HDL) cholesterol levels at all doses. Furthermore, there was a significant increase in low-density lipoprotein (LDL) cholesterol levels at the 40-mg and 80-mg doses.

TABLE 3. Safety Markers

				Oxandrolose	
		Placebo	20 mg	40 mg	80 mg
IIIV RNA by PCR (µL)	Baseline	204,915 ± 581,530	154,294 ± 377,496	156,563 ± 376,248	123,994 ± 391,468
	change week 12	-110,419 ± 654,209	1338 ± 734,313	-81,152 ± 297,702	-83,976 ± 346,641
CD4° T lymphocyte count (%)	Baseline	15.5 ± 10.1	14.9 ± 11.2	16.8 ± 11.4	15.1 ± 9.9
	change week 12	0.2 ± 4.1	1.4 ± 3.9	1.1 ± 5.3	0.8 ± 3.1
Hemoglobin (g/L)	Baseline	137 ± 20	137 ± 20	134 ± 19	137 ± 16
	change week 12	5 ± 15	-4 ± 13	-2 ± 14	-3 ± 15
Platelets (10°/L)	Baseline	221 ± 90.0	217 ± 67.2	228 ± 77.6	235 ± 70.6
,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,,	change week 12	3.1 ± 53.9	49.6 ± 86.7*	51.3 ± 103*	64.9 ± 59.1°
Creatinine (µmol/L)	Baseline	79.6 ± 17.7	79.6 ± 17.7	70.7 ± 17.7	79.6 ± 26.5
.,,	change week 12	0.0 ± 17.7	8.8 ± 26.5°	17.7 ± 26.5*	17.7 ± 17.7*
Creatinine kinase (U/L)	Baseline	192 ± 393	138 ± 220	114 ± 81	167 ± 217
	change week 12	-61 ± 424	58 ± 94*	88 ± 127*	70 ± 234°
AST (U/L)	Baseline	42.4 ± 24.7	39.9 ± 22.0	36.6 ± 20.8	39.5 ± 22.4
	change week 12	-2.6 ± 37.0	3.6 ± 23.8	12.1 ± 56.7	20.3 ± 38.3*
ALT (U/L)	Baseline	39.4 ± 27.2	42.7 ± 33.1	37.0 ± 28.5	40.1 ± 30.4
	change week 12	-1.6 ± 39.3	4.4 ± 38.3	19.2 ± 56.3°	37.5 ± 61.0°
Glucose (mmoi/L)	Baseline	5.2 ± 1.5	5.1 ± 1.1	4.9 ± 1.0	5.1 ± 0.9
, , , , , , , , , , , , , , , , , , , ,	change week 12	0.1 ± 1.0	0.1 ± 1.8	0.6 ± 2.4	0.1 ± 1.3*
Uric scid (umol/L)	Baseline	345 ± 95	339 ± 83	357 ± 107	333 ± 71
(,	change week 12	-12 ± 71	-54 ± 71*	-54 ± 101*	-77 ± 59*
Triglycerides (mmol/L)	Baseline	2.78 ± 2.99	3.99 ± 8.44	3.65 ± 4.90	2.28 ± 1.89
	change week 12	0.09 ± 2.01	-1.33 ± 7.10	-0.63 ± 2.96	0.12 ± 1.10
Chulesterol (mmol/L)	Bascline	4.6 ± 1.6	4.9 ± 2.9	4.6 ± 1.5	4.6 ± 1.3
, , , , , , , , , , , , , , , , , , , ,	change week 12	0.01 ± 1.0	-0.3 ± 2.4	0.4 ± 1.6	0.3 ± 1.2
LDL (mmol/L)	Baseline	2.8 ± 1.3	2.7 ± 1.1	2.6 ± 0.8	2.6 ± 0.8
	change week 12	0.1 ± 1.0	0.4 ± 1.3	0.7 ± 1.3*	0.8 ± 1.1*
HDL (mmol/L)	Baseline	1.0 ± 0.4	1.0 ± 0.4	0.9 ± 0.3	1.0 ± 0.4
	change week 12	-0.03 ± 0.3	-0.3 ± 0.3*	-0.3 ± 0.3*	-0.5 ± 0.4*
Lp(a) (mmol/L)	Baseline	0.7 ± 0.7	0.9 ± 1.0	0.5 ± 0.5	0.7 ± 0.8
	change week 12	0.2 ± 0.7	-0.2 ± 0.4	-0.2 ± 0.5*	$-0.6 \pm 0.7^{\circ}$

Values are mean ± SD.

*P < 0.017 vs. placebo.
Lota) indicates lipoprotein (a).

TARLE 4. Grade III or IV Toxicities and Reasons for Discontinuation

		Oxandrolone			
	Placebo	20 mg	40 mg	80 mg	Total
Grade III or IV toxicities					
AST	2	2	6	90	19
ALT	1	3	71	91	205
Total bilirubin	1	0	0	ı	2
LDH	0	0	0	t	1
Uric acid	0	0	0	t	1
Total grade III or IV toxicities	4	5	13	21	43
Reasons for discontinuation					
Adverse experience or abnormal laboratory tests	3	t	7	9	20
Noncompliance with protocol requirements	4	6	6	2	18
Voluntary patient withdrawal or requested removal	2	2	2	3	9
Patient moved or lost to follow-up	1	3	1	2	7
Death	1	3	- 1	1	6
Lack of efficacy	1	2	- 1	- 1	5
Intercurrent medical problem or disease and related complications	0	1	0	'	2
Total reasons for discontinuation	12	18	18	19	67

^{*}P = 0.054 vs. placebo

Six patents died during the placebo-controlled study (see Table 4), and 3 more died during the open-label phase or within 30 days of last receiving study medication during the placebo-controlled phase. Of the 9 subjects who died, 2 were on placebo, 3 were on 20 mg of oxandrolone, 2 were on 40 mg of oxandrolone, and 2 were on 80 mg of oxandrolone. There were no significant differences between the treatment groups in the numbers of infections, serious adverse events (SAEs). or milder adverse events. Seven SAEs were reported in 6 subjects on placebo, 20 SAEs were reported in 13 subjects on 20 mg of oxandrolone, 24 SAEs were reported in 14 subjects on 40 mg of oxandrolone, and 20 SAEs were reported in 14 subjects on 80 mg of oxandrolone. One hundred eighty-one different types of infections and adverse events were reported.

Overall dropout rates were similar among treatment groups (see Table 4). In some subjects, treatment discontinuation was prompted by more than I reason. There was a trend toward increased dropout because of an adverse experience or abnormal laboratory test results in the 40-mg and 80-mg oxandrolone groups attributable to treatment discontinuation for WHO grade III and IV elevations in AST and ALT.

Gonadal-Pituitary Function

Baseline total testosterone levels averaged close to the lower limits of normal (270 ng/dL; Table 5). At 12 weeks,

serum LH and FSH concentrations decreased significantly from baseline in all oxandrolone-treated groups, consistent with an androgenic action. Serum SHBG concentrations also decreased with increasing doses of oxandrolone, which also suggests an androgenic effect of oxandrolone (SHBG was determined in a subset of patients, and total testosterone levels in the subset were similar to those in the larger cohort; data not shown).

Total and free testosterone concentrations measured by direct RIA did not show a dose-related change. We used celite chromatography to separate testosterone from oxandrolone before RIA and found that serum total testosterone concentrations were significantly decreased from baseline at all doses of exandrolone but not with placebo treatment (see Table 5).

Open-Label Study

After the double-blind placebo-controlled study, a subset of subjects opted to take 20 mg of oxandrolone in an open-label study. All 4 groups receiving 20 mg of oxandrolone during this 12-week open-label phase continued to gain weight (Table 6). By the end of the open-label phase, there were no significant differences in weight gain among the groups. AST levels decreased; although AST levels remained above baseline, they were no longer significantly different from baseline (see Table 6).

DISCUSSION

Oxandrolone treatment was associated with significantly greater body weight gain above baseline than with placebo. A major portion of this weight gain occurred in the lean body compartment, as reflected in the significant gains in BCM, intracellular water, and serum creatinine levels. The gains in body weight during the double-blind phase of the study were sustained during the open-label phase of the study.

Oxandrolone administration has been shown to increase muscle protein synthesis in emaciated burn patients, 56 muscle mass and maximal voluntary strength in older men at risk for sarcopenia, 37-60 and weight in patients with cancer cachexia. Most previous studies have included small numbers of subjects, however; this study is the largest randomized placebo-controlled trial of an androgen in patients with HIVassociated weight loss.

Serum LH and FSH levels decreased significantly during oxandrolone administration, consistent with its androgenic activity. Whereas conventional measurement of testosterone did not show consistent decreases, assay after chromatographic separation did show suppression of testosterone, confirming the androgenic effect and indicating that oxandrolone or a metabolite cross-reacted in the conventional testosterone assay. This dose-ranging study did not include women; therefore, we cannot determine whether the level of androgenic activity seen with oxandrolone would have the expected detrimental virilizing effects in women.

Oxandrolone administration was generally well tolerated. Grade III and IV elevations of transaminases were observed in >5% of study participants, however, especially at the 80-mg dose. Careful monitoring of these parameters is therefore

¹P = 0.021 vs. placebo. 8Dose trend. P = 0.0047.

TABLE 5. Effect of Oxandrolone on Serum LH, FSH, Total and Free Testosterone, and SHBG Levels (baseline to week 12)

	•	Oxandrolone				
	Placebo	20 mg	40 mg	HO mg		
Testosterone by RIA (nmol/L)						
Daseline	7.7 ± 3.4	9.5 ± 4.6	9.2 ± 5.0	12.0 ± 13.69		
change week 12	2.6 ± 10.0	-1.7 ± 4.3	1.1 ± 10.0	-2.2 ± 5.6		
P	0.0581	0.0120	0.4523	0.012		
4	54	46	49	46		
Free testosterone (pmol/L)						
Baseline	101 ± 52	118 ± 59	114 ± 52	146 ± 139*		
change week 12	26 ± 109	-41 ± 55	-24 ± 86	-45 ± 51.3		
P	0.0838	0.0001	0.0581	0.0001		
9	54	46	49	46		
LH (U/L)						
Baseline	3.52 ± 2.61	4.07 ± 4.00	3.97 ± 2.66	4.03 ± 3.05		
change week 12	0.93 ± 4.68	-1.08 ± 2.33	-1.35 ± 2.90	-2.18 ±2.74		
P	0.1519	0.0029	0.0020	0.0001		
0	54	46	49	46		
FSH (U/L)						
Baseline	5.75 ± 4.95	5.12 ± 3.81	6.08 ± 4.52	4.81 ± 4.04		
change week 12	0.78 ± 2.94	-0.67 ± 3.37	-0.80 ± 3.08	-1.28 ± 2.06		
P	0.0563	0.1809	0.0753	0.0001		
n.	54	46	49	46		
SHBG (nmol/L)						
Baseline	44.8 ± 22.7	41.4 ±20.2	42.3 ± 23.4	44.2 ± 20.9		
change week 12	0.43 ± 16.0	-24.4 ± 21.3	-26.8 ± 23.1	-35.0 ± 17.2		
P	0.8751	1000.0	0.0001	0.0001		
9	35	23	31	24		
Testosterone by extraction and chri	omatography (ng/dL)					
Baseline	289 ± 153	269 ± 120	282 ± 117	314 ± 111		
change week 12	-1.5 + 156	-124 + 129	-126 ± 126	-209 ± 122		
P	0.504	0.001	0.001	0.001		
9	30	32	30	33		

indicated after the initiation of oxandrolone therapy. Furthermore, LDL levels increased and HDL levels decreased.

There has been considerable debate about what magnitude of change in body weight is clinically meaningful. An AIDS Clinical Trial Group (ACTG) expert panel on HIVassociated wasting expressed the opinion that a gain of 1.5 kg is clinically meaningful (Fred Sattler, MD, personal communication). The average weight gain at each of the oxandrolone doses exceeded 1.5 kg, whereas the increase in the placebo group was less than 1.5 kg. Only the 40-mg dose of oxandrolone induced more than a 1.5-kg increase in weight over that attained with placebo (an increase over placebo of 1.7 kg based on a 2.8-kg increase for 40 mg of oxandrolone vs. a 1.1-kg increase for placebo). For subjects whose BMI was ≤22.5 kg/m², all 3 doses of oxandrolone induced more than a 1.5-kg increase over placebo (20 mg induced a 1.9-kg increase, 40 mg induced a 2.1-kg increase, and 80 mg induced a 1.7-kg increase). In subjects whose BMI was ≤22.5 kg/m², the mean increases in BCM in patients treated with the 40- or 80-mg dose of exandrolone were also greater than 1.5 kg above that attained with placebo. These changes in weight and

BCM compare favorably with those observed during administration of rhGH^{27,28} and testosterone. ³¹⁻³⁶ In a meta-analysis of placebo-controlled, randomized, clinical trials of testosterone, the average gain in lean body mass was 1.3 kg in testosterone-treated HIV-infected men. ⁶¹

In spite of significant body weight gains and lean mass accretion, total work output during treadmill exercise did not significantly change during treatment. This is consistent with the growing body of data that androgenic steroids increase muscle mass but do not affect measures of endurance, such as treadmill performance. 82-64 Reports of randomized clinical trials published subsequent to the initiation of this study have reported significant gains in maximal voluntary strength with androgen supplementation of HIV-infected men with weight loss³⁵; gains in muscle strength are generally proportional to increases in muscle mass.³⁵

Participants in this study were able to consume a wellbalanced diet at study entry as assessed by a dietitian. In developing countries of Africa and Asia, many HIV-infected patients have an overall energy deficit, with varying macroand microputrient deficiencies. We do not know whether

TABLE 6. Change From Baseline in Weight, AST, and ALT in Subjects Continuing in the Open-Label (20 mg) Study

		Oxandrolone			
	Placebo	20 mg	40 mg	80 mg	
Weight gain (kg)					
n	53	46	48	46	
Double-blind placebo phase				2.5 ± 3.0	
0 to 12 weeks	1.3 ± 3.0	2.0 ± 3.9	3.0 ± 3.3°	2.3 £ 3.0	
Open-label 20-mg phase					
12 to 24 weeks	1.6 ± 4.4	1.4 ± 2.5	0.3 ± 2.7	1.1 ± 3.0	
Liver function tests					
n	42	40	39	38	
AST					
Baseline to 24 weeks	6.2 ± 45.4	-1.4 ± 22.1	6.8 ± 30.2	18.4 ± 39.9	
ALT					
Baseline to 24 weeks	7.6 ± 43.9	7.6 ± 69.3	17.4 ± 42.1	35.1 ± 70.8	
*P < 0.004 vs. placebo.					

androgen administration would be efficacious in preventing weight loss in HIV-infected patients with severe wasting or in nutritionally depleted individuals.

Administration of exandrolone has been associated with significant decreases in plasma HDL cholesterol levels and increases in LDL cholesterol levels. The administration of the 40- and 80-mg doses was associated with significant increases in ALT and AST; these increases were transient and returned toward baseline in most subjects. Treatment discontinuations attributable to persistent and marked increases in transaminases were common and occurred in more than 5% of individuals. We found no increase in bilirubin or alkaline phosphatase.

The gains in body weight and BCM were related to oxandrolone dose. Similarly, there were dose-dependent increases in AST and ALT levels and common treatment discontinuations attributable to AST and ALT elevations. Thus, the best trade-off between the anabolic effects and AST and ALT elevation was achieved at the 40-mg daily dose. The therapeutic efficacy and safety of this dose should be further evaluated in subsequent clinical trials.

The decreases in HDL and increases in LDL represent a proatherogenic lipoprotein profile. Clinicians therefore need to weigh the risk-benefit ratio of this therapy. Wasting syndrome predicts a significant risk of complications and death, but even studies as large as this one are not large enough and have not been carried out long enough to determine whether reversal of that risk occurs with treatment of wasting and to determine the risk of cardiovascular disease. The risk of atherosclerosis predicted by this lipoprotein profile suggests that such therapy should be restricted to those with significant wasting or should be terminated when wasting has improved. Mean CD4 lymphocyte counts in this study were >200 × 10⁶/L, which is higher than in most earlier studies of HIV-associated wasting (which often had mean values ≤50 × 106/L), indicating better health later in the epidemic. In that light, future studies should likely exclude those with obesity even in the presence of weight loss. In post hoc analysis, we found that the 20-mg dose was more

effective in those with a BMI at entry of ≤22.5 kg/m². The lower dose was accompanied by lesser increases in LDL and transaminases. Thus, a prospective study excluding obese patients could establish that a 20-mg dose is efficacious and associated with a lower frequency of adverse events.

A number of therapies, including dronabinol, mege-strol acetate.^{24,25} and rhGH, ²⁶⁻²⁹ are approved for the treatment of HIV-associated wasting. Orexigenic agents, such as dronabinol and megestrol acetate, increase appetite but have not been shown to increase lean body mass. rhGH was approved for treatment of patients with HIV-associated was approved for a trial that demonstrated increases in fatfree mass and increased performance on treadmill testing. rhGH is expensive, however, and its administration is associated with adverse effects at the approved doses. Oxandrolone compares favorably with rhGH in terms of the weight and BCM gain as well as retail cost. Furthermore, oxandrolone did not reduce fat stores and is associated with a lower frequency of adverse events than rhGH. Therefore, it may be viewed as an adjunct or alternative to rhGH for the treatment of patients with HIV-associated weight loss. Of 2 recent smaller studies on the use of nandrolone, an injectable anabolic steroid, for AIDS wasting, one reported that nandrolone induced a similar gain in weight⁶⁷ to the increase seen here with oxandrolone, whereas the other found that nandrolone induced a larger gain in weight than we report with oxandrolone. Oxandrolone has the advantage of oral administration, however, which may be important in patients with loss of muscle and fat, such as occurs in AIDS wasting. Further studies are needed to determine the efficacy of oxandrolone in improving muscle strength, physical function, and health-related QOL in HIV-infected patients with weight loss.

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